Commission Directive 2008/78/EC of 25 July 2008 amending Directive 98/8/EC of the European Parliament and of the Council to include propiconazole as an active substance in Annex I thereto (Text with EEA relevance) (repealed)

COMMISSION DIRECTIVE 2008/78/EC

of 25 July 2008

amending Directive 98/8/EC of the European Parliament and of the Council to include propiconazole as an active substance in Annex I thereto

(Text with EEA relevance) (repealed)

THE COMMISSION OF THE EUROPEAN COMMUNITIES,

Having regard to the Treaty establishing the European Community,

Having regard to Directive 98/8/EC of the European Parliament and of the Council of 16 February 1998 concerning the placing of biocidal products on the market⁽¹⁾, and in particular the second subparagraph of Article 16(2) thereof,

Whereas:

- (1) Commission Regulation (EC) No 1451/2007 of 4 December 2007 on the second phase of the 10-year work programme referred to in Article 16(2) of Directive 98/8/EC of the European Parliament and of the Council concerning the placing of biocidal products on the market⁽²⁾ establishes a list of active substances to be assessed, with a view to their possible inclusion in Annex I, IA or IB to Directive 98/8/EC. That list includes propiconazole.
- (2) Pursuant to Regulation (EC) No 1451/2007, propiconazole has been evaluated in accordance with Article 11(2) of Directive 98/8/EC for use in product type 8, wood preservatives, as defined in Annex V to Directive 98/8/EC.
- (3) Finland was designated as Rapporteur Member State and submitted the competent authority report, together with a recommendation, to the Commission on 5 April 2006 in accordance with Article 14(4) and (6) of Regulation (EC) No 1451/2007.
- (4) The competent authority report was reviewed by the Member States and the Commission. In accordance with Article 15(4) of Regulation (EC) No 1451/2007, the findings of the review were incorporated, within the Standing Committee on Biocidal Products on 29 November 2007, in an assessment report.
- (5) The review of propiconazole did not reveal any open questions or concerns to be addressed by the Scientific Committee on Health and Environmental Risks.
- (6) It appears from the examinations made that biocidal products used as wood preservatives and containing propiconazole may be expected to satisfy the requirements laid down in Article 5 of Directive 98/8/EC. It is therefore appropriate to include propiconazole in Annex I for product type 8, in order to ensure that in all Member

States authorisations for biocidal products used as wood preservatives and containing propiconazole can be granted, modified, or cancelled in accordance with Article 16(3) of Directive 98/8/EC. However, unacceptable risks were identified for the *in situ* treatment of wood outdoors and for treated wood exposed to weathering. Authorisation of these uses will require the submission of data demonstrating that the products can be used without unacceptable risks to the environment.

- (7) In the light of the findings of the assessment report, it is appropriate to require that products containing propiconazole and used as wood preservatives must be used with appropriate personal protective equipment, that risk mitigation measures are applied to protect the soil and aquatic compartments and that related instructions are provided, in accordance with Article 10(2)(i)(d) of Directive 98/8/EC.
- (8) It is important that the provisions of this Directive be applied simultaneously in all the Member States in order to ensure equal treatment of biocidal products on the market containing the active substance propiconazole and also to facilitate the proper operation of the biocidal products market in general.
- (9) A reasonable period should be allowed to elapse before an active substance is included in Annex I in order to permit Member States and the interested parties to prepare themselves to meet the new requirements entailed and to ensure that applicants who have prepared dossiers can benefit fully from the 10-year period of data protection, which, in accordance with Article 12(1)(c)(ii) of Directive 98/8/EC, starts from the date of inclusion.
- (10) After inclusion, Member States should be allowed a reasonable period to implement Article 16(3) of Directive 98/8/EC, and in particular, to grant, modify or cancel authorisations of biocidal products in product type 8 containing propiconazole to ensure that they comply with Directive 98/8/EC.
- (11) Directive 98/8/EC should therefore be amended accordingly.
- (12) The measures provided for in this Directive are in accordance with the opinion of the Standing Committee on Biocidal Products,

HAS ADOPTED THIS DIRECTIVE:

Article 1

Annex I to Directive 98/8/EC is amended in accordance with the Annex to this Directive.

Article 2

Transposition

1 Member States shall adopt and publish, by 31 March 2009 at the latest, the laws, regulations and administrative provisions necessary to comply with this Directive. They shall forthwith communicate to the Commission the text of those provisions and a correlation table between those provisions and this Directive.

They shall apply those provisions from 1 April 2010.

When Member States adopt those provisions, they shall contain a reference to this Directive or be accompanied by such a reference on the occasion of their official publication. Member States shall determine how such reference is to be made.

2 Member States shall communicate to the Commission the text of the main provisions of national law which they adopt in the field covered by this Directive.

Article 3

This Directive shall enter into force on the 20th day following its publication in the *Official Journal of the European Union*.

Article 4

This Directive is addressed to the Member States.

Done at Brussels, 25 July 2008.

For the Commission
Stavros DIMAS
Member of the Commission

ANNEX
The following entry 'No 8' is inserted in Annex I to Directive 98/8/EC:

No	Commoname	n IUPAC nameIde numbers	Minimulation of the active substant in the biocidal product as placed on the market	n of inclusion	for compliant with Article 16(3) (except for products containing more than one active substance for which the deadline to comply with Article 16(3) shall be the one set out in the last of the inclusion decisions relating to its active	ng ee,	Product type	Specific provisions ^a
·8	propicona	z io[∉2- (2,4-	930 g/kg	1 April 2010	substance 31 March	31 March	8	Member States
		dichlorop propyl-1,3 dioxolan-	3- 2-]-1H-1,2,4		2012	2020		shall ensure that authorisations are subject to the

a For the implementation of the common principles of Annex VI, the content and conclusions of assessment reports are available on the Commission website: http://ec.europa.eu/comm/environment/biocides/index.htm

ı	CAC	ſ	I	1	I		Callarri -
	CAS						following
	No:						conditions:
	60207-90	-1					In view
							of the
							assumptions
							made
							during
							the risk
							assessment,
							products
							authorised
							for
							industrial
							and/or
							professional
							use,
							must
							be used
							with
							appropriate
							personal
							protective
							equipment,
							unless it
							can be
							demonstrated
							in the
							application
							for
							product
							authorisation
							that
							risks to
							industrial
							and/or
							professional
							users can be
							reduced
							to an
							acceptable
							level by
							other
							means.
							In view
							of the
							risks
							identified
							for the
							soil and
							aquatic
							aquatio

a For the implementation of the common principles of Annex VI, the content and conclusions of assessment reports are available on the Commission website: http://ec.europa.eu/comm/environment/biocides/index.htm

				compartments
				appropriate
				risk
				mitigation
				measures
				must be
				taken to
				protect
				those
				compartments.
				In
				particular,
				labels
				and/or
				safety
				data
				sheets of
				products
				authorised
				for
				industrial
				use shall
				indicate
				that
				freshly
				treated
				timber
				must be
				stored
				after
				treatment
				under
				shelter
				or on
				impermeable
				hard
				standing
				to
				prevent
				direct
				losses
				to soil
				or water
				and that
				any
				losses
				must be
				collected
				for
				reuse or
				disposal.
				uisposai.

For the implementation of the common principles of Annex VI, the content and conclusions of assessment reports are available on the Commission website: http://ec.europa.eu/comm/environment/biocides/index.htm

								In addition, products cannot be authorised for the <i>in situ</i> treatment of wood outdoors or for wood that will be exposed to weathering unless data is submitted to demonstrate that the product will meet the requirements of Article 5 and Annex VI, if necessary by the application of appropriate risk mitigation measures.'
--	--	--	--	--	--	--	--	--

a For the implementation of the common principles of Annex VI, the content and conclusions of assessment reports are available on the Commission website: http://ec.europa.eu/comm/environment/biocides/index.htm

- (1) OJ L 123, 24.4.1998, p. 1. Directive as last amended by Directive 2008/31/EC (OJ L 81, 20.3.2008, p. 57).
- (2) OJ L 325, 11.12.2007, p. 3.